REPORT DOCUMENTATION PAGE AFRL-SR-BL-TR-98-Public reporting burden for this collection of information is estimated to average 1 hour per response, inc gathering and maintaining the data needed, and completing and reviewing the collection of information. collection of information, including suggestions for reducing this burden, to Washington Headquarters Separate Highway, Suite 1204, Arlington, VA 22202-4302, and to the Office of Management and Budget, Paperwork Reductions of the Company of the t of this efferson 1. AGENCY USE ONLY (Leave blank) 2. REPORT DATE 3. REPORT TYPE AND DATES COVERED May 15, 1998 Final Technical Report 01 Mar 95 to 28 Feb 98 4. TITLE AND SUBTITLE 5. FUNDING NUMBERS ORGANIC POLLUTANTS IN SOILS, AS STUDIED BY NUCLEARMAGNETIC F49620-95-1-0192 RESONANCE 6. AUTHOR(S) Gary E. Maciel Willard L. Lindsay 7. PERFORMING ORGANIZATION NAME(S) AND ADDRESS(ES) 8. PERFORMING ORGANIZATION REPORT NUMBER Department of Chemistry Colorado State University Fort Collins, CO 80523 9. SPONSORING/MONITORING AGENCY NAME(S) AND ADDRESS(ES) 10. SPONSORING/MONITORING AGENCY REPORT NUMBER . AFOSR/NA 110 Duncan Avenue, Ste B115 F49620-95-1-0192 Bolling AFB, DC 20332-8050 11. SUPPLEMENTARY NOTES 19980617 029 12a. DISTRIBUTION AVAILABILITY STATEMENT Approved for public release; distribution unlimited. 13. ABSTRACT (Maximum 200 words) This three-year project focussed on the development and application of state-of-the art NMR (nuclear magnetic resonance) techniques to elucidate the fundamental behaviors of certain organic pollutants (e.g., benzene, CCl, trichloroethylene, ethylene glycol) when adsorbed in typical soils, as represented in most of this study by the following major soil components: humics (humic acid, fulvic acid, humin), clays (montmorillonite, kaolinite) and silica. The first stage was the isolation/separation and detailed ¹³C NMR characterization of the humic materials from a soil of southwestern Colorado; this work is resulting in the most detailed chemical-structural elucidation ever carried out on such a three-component humic suite. ¹³C NMR studies revealed a spectacular case of co-contamination in reactions between CC1₄ and benzene adsorbed in montmorillonite clays, yielding such Friedel-Crafts products as benzophenone, benzoic acid and (C₆H₅)₃C⁺ (the last being analogous to the product in a "ship-in-a-bottle" reaction between the two initial components in HY zeolite). 13C NMR also elucidated the photo-assisted decomposition of trichloroethylene absorbed on soil components (forming C1₂CHCO₂H, Cl₃CCO₂H, Cl₃CCHCl₂, etc.), a much slower analog of the well -known conversions employing TiO₂(s) as the adsorbent. ²H NMR lineshape studies elucidated the detailed local motions of benzene, trichloroethylene and ethylene glycol adsorbed on major soil components. 14. SUBJECT TERMS 15. NUMBER OF PAGES 16. PRICE CODE

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FINAL TECHNICAL REPORT

ORGANIC POLLUTANTS IN SOILS, AS STUDIED BY NUCLEAR MAGNETIC RESONANCE

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Objectives

The goal of this project was to elucidate as much detail as possible regarding the chemical/physical behaviors of certain organic pollutants (e.g., benzene, ethylene glycol, trichloroethylene and fluorocarbon ethers) in soils, initially in terms of primary soil constituents, i.e., humic acids, fulvic acids and clays (kaolinite and montmorillonite). This study was designed to provide the kind of fundamental chemical/physical information that can be used by others for constructing models of the sequestering, decomposition and transport of organic pollutants and their residues in soil/groundwater systems. Because of time constraints, work on the fluorocarbon ethers was only minimal, and will not be discussed in this report.

Organization of the Effort

The primary tool of this study was nuclear magnetic resonance (NMR), employing detection of primarily the nuclides, ¹H, ²H, ¹³C, ¹⁹F, ²⁷Al and ²⁹Si. Both liquid solutions (e.g., pollutant + humic or fulvic acid) and solid samples (e.g., pollutant/clay or pollutant/humic "complex") were studied by high-resolution NMR techniques. Emphasis was heavily on the latter. Magnetic fields ranging from 2.1 T (90 MHz proton resonance frequency) to 14 T (600 MHz proton frequency) were employed.

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This report is organized primarily on the basis of providing a relatively detailed description of each major aspect of the project. In several cases, the data resulted in manuscripts that have been published, accepted or submitted for publication, or are at the stage of a draft of a manuscript being revised to submit for publication. In most of those cases, copies of the manuscripts are included as part of this report, and the work is described only briefly elsewhere in the report; in cases where a manuscript is in preparation, but is not quite ready at this time, the manuscript(s) will be provided upon request.

This project focused on the development and application of multinuclear NMR strategies for elucidating the fundamental physical/chemical (*abiotic*) behaviors (chemical reactions, interactions, local mobilities) of specific organic pollutants in major soil components (clays, silica and humic materials – humic acid, fulvic acid, humin) and, to a lesser degree, whole soils. This overall effort, and this report, can be viewed in terms of the following separate, but related, theme areas: 1) choosing and characterizing the clays to be used for pollutant/clay studies; 2) isolation and characterization of humic materials for pollutant studies; 3) local mobilities of certain organic pollutants adsorbed on/in soil components; 4) photoassisted decomposition of Cl₂C=CHCl adsorbed on soil components; 5) cocontamination of zeolites and clays with CCl₄ and benzene; and 6) the silica surface.

Accomplishments/New Findings

1. Clays. One of the important tasks of the first several months of this project was selecting and characterizing suitable clay and humic materials on which to base the pollutant/soil system samples and studies. The clays that were ultimately selected, based on x-ray powder diffraction, elemental analysis and ²⁹Si and ²⁷Al NMR data, are a kaolinite and a Camontmorillonite obtained from the Missouri Clay Repository, Columbia, MO. Figures 1-3 show ²⁷Al and ²⁹Si NMR spectra of these clay samples, based on magic angle spinning (MAS) and, in the ²⁹Si spectra, either direct polarization (DP, generation of ²⁹Si spin polarization directly by ²⁹Si spinlattice relaxation) or cross polarization (CP, a transfer of spin polarization from protons to ²⁹Si nuclei, based on the existence of strong, static components of ¹H-²⁹Si magnetic dipole-dipole interactions).

In the ²⁷Al NMR spectra shown in Figure 1, one sees the expected dominance of the octahedral (Oh) peak at about 0 ppm and a much smaller tetrahedral (T_d) aluminum peak at about 50-60 ppm, the latter representing a small, but significant quantity of Al-for-Si substitutions in the tetrahedral (nominally silicon) planes of the clay structures. In the ²⁹Si NMR spectra of Ca-montmorillonite (Fig. 2) and kaolinite (Fig. 3), one sees the strong tetrahedral silicon peak expected for clay structures. For the Camontmorillonite, there is also a small peak at about -110 ppm, due to silica. The fact that this peak shows up so strongly in the CP spectrum, where the observed ²⁹Si spin polarization has been transferred from protons (presumably in either surface SiOH groups or physisorbed H₂O), implies a higher surface areas than one would expect for the quartz-like silica normally assumed to be present in soils. The fact that metal ion exchange with Cu²⁺ enhances the relative intensity of the silica peak of Camontmorillonite (Fig. 4), along with broadening both peaks (as expected for a paramagnetic agent), indicates that the added Cu²⁺ is incorporated into, or complexed with, both the clay and the silica; this observation may represent an important step in elucidating the true nature of this silica.

The humic samples on which much of this project is being based were isolated from a soil that we collected in the Uncompangre National Forest of southwestern Colorado. The classical soil separation procedures yielded humic acid, fulvic acid and humin fractions, for which the ¹³C CP-MAS NMR spectra are shown in Figure 5, which also provides structural assignments for the major peaks and shoulders.

2. Humic Materials. This aspect of the project, which we had expected be completed shortly after the first year of the project, has turned out to be a much bigger, and much more important, task than we had envisioned. This work is now nearly complete and a full paper on it will be submitted shortly. After we learned that commercially available humic materials are either not available in sufficient quantities to support our program (e.g., samples from the International Humic Substances Society) or of questionable origin and significance (e.g., from Aldrich), we decided to collect our own soil and perform the separations into humic acid, fulvic acid and humin, based on the classical acid/base dissolution procedures. This approach also offered the prospect of obtaining the three types of samples from *one* soil source, so the suite of three materials will have maximum significance to soil scientists and organic geochemists. Since we were choosing the soil ourselves, we were free to choose one that has a low-iron

content, minimizing the NMR complications that one encounters from paramagnetic and ferromagnetic iron sites in a soil. We collected a soil from the Uncompandere National Forrest of southwestern Colorado, about 18 inches below the "leaf litter." The "classical" (wet chemical) characteristics were also measured for this soil.

The ¹³C CP-MAS spectra of the whole soil and the three solid organic components that we separated from it are shown in Figure 5. Shown with the spectra are peak assignments based on chemical shifts of known chemical functionalities and on spectral editing (vide infra). One sees a wide variety of organic chemical functionalities in these humics, making them ideal for the pollutant/soil interaction studies of this project. One notes the close match-up between the computer-synthesized spectrum of Fig. 1e (a linear combination of the spectra of humic acid, fulvic acid and humin) and the spectrum of the whole soil, indicating that any chemical changes that occurred in the chemical structures of the organic soil components during the acid/base separation procedure are minimal or subtle. Figure 6 shows the liquid-sample ¹³C NMR spectrum of the fulvic acid component. Its improved resolution, relative to that of the CP-MAS spectrum of Fig. 1d, is due primarily to the averaging by liquid-state motion of small differences in isotropic chemical shifts (due to corresponding small differences in local geometry, e.g., conformation) for each particular class of functional group.

In order to identify, or confirm, the chemical structure(s) responsible for each spectral region of the spectra shown in Figure 5, spectral editing experiments were carried out. The most primitive of these techniques is the popular "interrupted-decoupling" or "dipolar-dephasing" technique, which differentiates between -CH₂- or C-H groups and -CH₃ or CH₀ groups in terms of their behaviors under a period during which the ¹H decoupler is turned off. From the dipolar-dephasing spectra on the three humic samples, shown in Figure 7, one can make the structural distinctions indicated above.

A higher level of spectral editing, distinguishing among the four carbon types, CH₃, CH₂, CH and CH₀ (no directly attached hydrogens) is achievable, at great cost in spectrometer time, by the combination of CP experiments, including cross depolarization, developed by Zilm and Wu and co-workers. These techniques have been applied to the three humic materials of this study. Figure 8 shows the individual component spectra, derived after the appropriate combinations of spectra obtained according to the published approach, and the composite spectra obtained by addition of

these components. Figure 9 shows the general procedure used for spectral editing.

In addition to applying published spectral editing methods, we attempted unsuccessfully to develop an alternate method, based on increasing T_{2C} sufficiently (at lower temperatures) to permit J_{CH} -based spectral editing, in direct analogy to what is done routinely on liquid samples. Although continued work on the structural characterization of humic materials is not now a major issue in a renewed project, we plan to see if an improved ^{13}C spectral-editing technique can be based on a synchronized combination of MAS with a suitable multiple-pulse sequence.

In order to quantitate the intensities of the various spectral regions assigned by the combination of chemical shift recognition and spectral editing, we carried out spin counting experiments on the three humic samples of this study. For this purpose, DP-MAS experiments (direct polarization, no CP) were carried out, based on long repetition delays (4s) and relaxation corrections computed from ¹³C T₁ values that we had determined. Quantitation in corresponding CP-MAS experiments was achieved by variable contact-time determinations (Figures 10-12) of the CP-relevant relaxation parameters. In the quantitation experiments, a previously developed ¹³C-labeled bicyclic ketone (labeled carbonyl peak outside the main spectral region of these humic samples) was contained in a small capillary as an intensity reference. Figure 13 shows typical high-S/N, CP-MAS and DP-MAS spectra obtained for this purpose, showing the T_{1C} values and relative populations for each indicated region of the spectrum.

¹³C CP-MAS and DP-MAS spectra we have obtained on the Uncompahgre humic samples (e.g., Figure 13) have unusually and remarkably high quality. We do not believe that spectra of this quality have been obtained by any other laboratory on humic samples. With such high quality (e.g., high S/N), it is possible to extract substantial structural detail via spectral deconvolution. Figure 14 shows pictorially the results of one such analysis.

The spectral editing and spin counting experiments described here are now being completed on the humin sample. These data are being interpreted in terms of hypothetical structural models. Before submitting the paper on this study (probably about June, 1998), we will have elucidated more detailed structural knowledge (both qualitative and quantitative) about the

organic components of Uncompangre soil than has ever been available previously on *any* soil.

3. Mobilities of Adsorbed Organic Pollutants. Another productive strategy of this project was the application of ²H NMR lineshapes of deuterium-labeled organics adsorbed on/in major soil components. To various levels of detail and completeness, a series of pollutant species were examined in this manner. These include HOCD₂CD₂OH, benzene (C₆D₆) and Cl₂C=CDCl. The overall strategy was to obtain ²H lineshapes, which are dominated by deuterium's orientation-dependent nuclear electric quadrupole interaction, as a function of temperature, and then try to model each line shape with computer-simulated line shapes based on specific motional models for the C-D moiety. From these comparisons, one can usually elucidate the detailed nature of motion at each temperature (from which an activation energy can also be obtained).

Our most detailed ²H lineshape study has been on ethylene glycol (used heavily in aircraft de-icing), for which a detailed paper is in preparation, describing the dependence of the motion of ethylene glycol adsorbed on/in a Ca-montmorillonite clay as a function of loading level, water content and metal-ion exchange. Figure 15 shows a representative set of ²H spectra measured from -125° C to 30° C, which show a general narrowing and change in line shape as the sample temperature was increased. Figure 16 shows the computer-generated lineshape simulations that correspond to the experimental spectra of Fig. 15; these simulations were based on a simulation model in which the C-D moieties are exchanging between two populations. In one population the C-D bond reorients (essentially isotropically) among the positions corresponding to the twelve vertices of an icosohedron; in the other population, the C-D moieties occupy (randomly) just one orientation. The populations and exchange rates of these simulations are listed in Figure 16.

An analogous analysis has been carried out for ethylene glycol adsorbed on humic acid (Figure 17) and on a whole soil. Manuscripts based on these results are currently being prepared for publication.

²H NMR lineshape studies were carried out on benzene adsorbed on Ca-montmorillonite at different loading levels and clay-moisture contents. Figures 18 and 19 display representative spectra. Simulations (not given here) show that the benzene molecules in these systems manifest, at

temperatures above the melting point of benzene, higher levels of local mobility than seen for samples with polar functional groups. These results have been described in a manuscript submitted for publication.

A few ²H NMR lineshape experiments have also been carried out on Cl₂C=CDCl adsorbed on Ca-montmorillonite. Figure 20 shows representative spectra. These results have been included in a manuscript submitted for publication.

4. Photo-Assisted Decomposition of Trichloroethylene Adsorbed on Soil Components. This work has been described in detail in a manuscript that has been submitted for publication. Only a few highlights are described here.

The general focus in this specific theme area was to begin characterizing the photo-induced chemistry that occurs with chlorohydrocarbons at the air/soil interface. This focus is in contrast to the huge literature that already exists on the photo-induced decomposition of chlorohydrocarbons in model reactors designed for the clean-up of waste water. Apparently the most successful photochemistry-based remediation systems rely on $TiO_2(s)$ as a substrate. Our work was directed to substrates that are highly abundant in soils, e.g., (especially) clays, silica and humics.

Our ¹³C NMR-based study has shown that trichloroethylene adsorbed on a variety of substrates (clay, silica, humic acid) decomposes to a variety of products (e.g., Cl₂CHCO₂H, Cl₃CCO₂H, Cl₃CCHCl₂, OCCl₂ and CO₂) under the influence of long-wavelength UV radiation, over a period of a few hours to several days. Figure 21 shows the ¹³C MAS spectra obtained on Cl₂C=CHCl/Ca-montmorillonite samples subjected to near-UV irradiation over periods of up to 20 days. As indicated on the figure, all of the abovementioned decomposition products can be seen in the ¹³C DP-MAS spectra, except the most volatile components, CO₂ and Cl₂CO (phosgene), which escape the reaction cell. After 10-20 days, most of the components are gone from the spectrum, largely decomposed to gaseous products (which we quantitated by using suitable collection traps). Figure 22 shows that this behavior is not specific to Ca-montmorillonite; analogous behaviors, albeit different in detail, are observed with a variety of other soil-based substrates (i.e., ion-exchanged montmorillonites, kaolinite, silica, humic acid and soil).

At this stage we have considerable detail on the photochemically-assisted product formation and product distribution for Cl₂C=CHCl/Ca-

montmorillonite, almost no such details for Cl₂C=CHCl adsorbed on any other substrate, and no information on the corresponding photophysics of any of these processes. These are areas that we propose to research as a major part of a subsequent project.

5. Co-Contamination with Benzene and CCl₄ Adsorbed on Clays and Zeolites. This aspect of our project has been described in two publications. Therefore, again, only highlights are described here.

Our first hint of "co-contaminant chemistry" came with the observation that $(C_6H_5)_3C^+$ carbenium ions could be generated inside the cavity of HY zeolite, from a mixture of benzene and CCl₄. These carbocations are presumably formed by chemical processes of the Friedel-Crafts variety inside the zeolite cavities in a kind of "ship in a bottle" synthesis (Fig. 23). That the carbocations were formed *inside* the zeolite, rather than on the outer surfaces of the zeolite particles, was demonstrated by chemical responses of the "trapped" $(C_6H_5)_3C^+$ species to reagents of various diagnostic sizes. Figure 24 shows the ¹³C NMR evidence for the formation of the $(C_6H_5)_3C^+$ ions. Supporting evidence on the motional characteristics of (C₆H₅)₃C⁺ cations inside the zeolite cavities, or adsorbed on the outside particle surfaces, was provided by ²H NMR results. Figure 25 shows representative ${}^{2}H$ NMR lineshapes for a sample containing $(C_{6}H_{5})_{3}C^{+}$ inside zeolite cavities; computer simulations of the ²H lineshape (not given here) show that this species executes a type of motion that is analogous to that of a constrained, but isotropically reorienting, species.

Based on the provocative results described briefly above on the Friedel-Crafts chemistry occurring in HY zeolite between two important pollutants (C_6H_6 and CCl_4), and recognizing the catalytic similarities between certain zeolites and certain clays, we examined the CCl_4/C_6H_6 system in clay samples. Figure 26 shows ¹³C CP-MAS and DP-MAS spectra of CCl_4 , benzene and the $CCl_4-C_6H_6$ combination adsorbed on Camontmorillonite, Zn^{2+} -exchanged montmorillonite, kaolinite, Zn^{2+} -exchanged kaolinite and K-10 montmorillonite. One sees in Fig. 26 that, especially for the Zn^{2+} -exchanged montmorillonites, chemistry of the Friedel-Crafts type occurs in these clay systems. In these cases, in contrast, to the zeolite analog, other Friedel-Crafts products are formed, e.g., $C_6H_5CO_2H$ and $(C_6H_5)_2CO$.

6. The Silica Surface. A long-term theme in our research program has been the detailed characterization of structure, dynamics and chemical

reactivity at silica surfaces. Since these issues are *a priori* highly relevant to the subject of pollutant behaviors in soils, some attention has been directed in this project to NMR studies that focus on various characteristics of silica surfaces, especially the distribution and structural/ dynamical characteristics of silanols (SiOH groups) on the surface. In the early stages of this project we completed a detailed ¹H and ²⁹Si NMR study of fumed silicas, focusing on the differences and similarities between the fumed silica surface and that of silica gel. This work was described in detail in a paper published in *J. Am. Chem. Soc.*

In our latest publication on the silica surface (based, at least in part, on this project), we presented a generalized version of a previously suggested model of the silica surface, in which geminal silanols are situated on surface segments similar to (100)-type faces of the β-cristobalite structure and single silanols are situated on surface segments similar to corresponding (111)-type faces. This model was supported by an extensive body of spectroscopic data, especially our own ²⁹Si and ¹H NMR data, gathered over a period of roughly 15 years. In this model single silanols on the same (111)-type surface segment cannot form hydrogen bonds with each other. Whether or not adjacent geminal silanols on the same (100)-type surface segment can form hydrogen bonds with each other depends on the relative orientation of their hydroxyl groups. When two surface segments of either (100)-type or (111)-type intersect convexly, hydroxyl groups cannot participate in hydrogen bonding across the intersection; but when two surface segments intersect concavely, those silanols situated at the intersection can form hydrogen bonds with their counterparts across the intersection. All the hydrogen-bonding silanols in this generalized β-cristobalite model have a common feature: when any two silanols are hydrogen bonded to each other, the two silicon atoms containing them are also situated on the same (100)type surface segment. The generalized β-cristobalite surface model can also explain the reversible dehydroxylation and rehydroxylation process on silica surfaces. Both single and geminal silanols participating in hydrogen bonding are most easily dehydroxylated under evacuation at temperatures between 170 and 450° C and form low-strain bicyclo[3.3.0]octasiloxane rings. The mode of dehydroxylation on a silica surface undergoes a transformation between 450 and 650° C, yielding highly strained trisiloxane rings for dehydroxylation at $T \ge 650^{\circ}$ C, according to this model. We expect this model to prove useful in interpreting data of this project on the interactions of organic pollutants with silica.

Personnel Supported

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Postdocs (partially supported): I.-S. Chuang, J. Xiong, J. Yang, H. Lock

Graduate Students: D. Keeler, T. Tao, C. Liu, M. Seger

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Ting Tao, Jane J. Yang and Gary E. Maciel, "Photo-Assisted Decomposition of Trichloroethylene on Clay," *Environ. Sci. Technol.*, submitted.

Jincheng Xiong, Ting Tao and Gary E. Maciel, "²H NMR Characterization of Motion and Mobility of Zeolite-Immobilized Triphenylmethyl Carbocations," *J. Phys. Chem.*, submitted.

Inventions/Patents

None

Acknowledgement/Disclaimer

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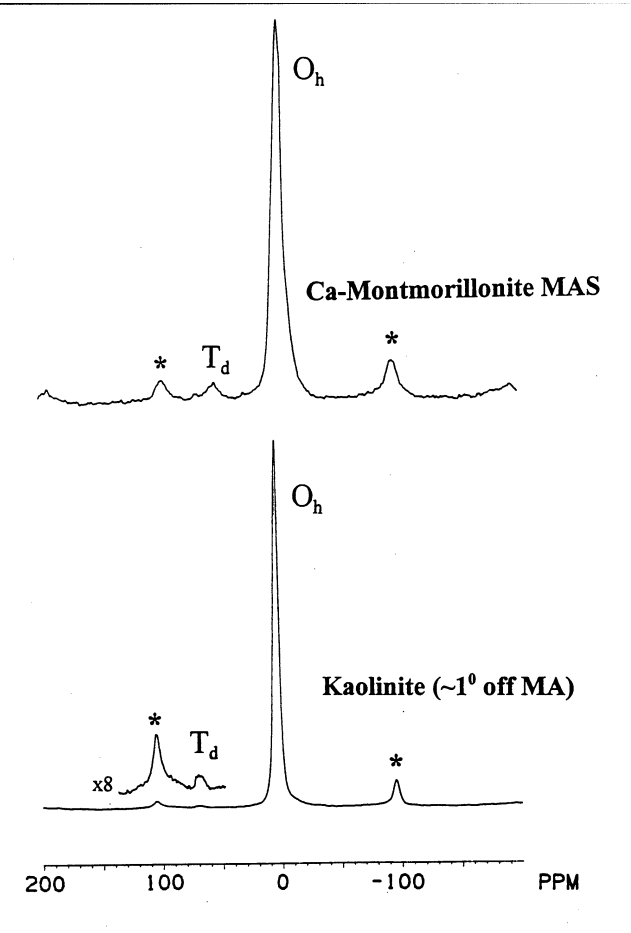


Figure 1. 27 Al MAS (magic-angle spinning) NMR spectra of clay minerals. Upper, Ca-montmorillonite. Bottom, kaolinite. T_d designates the tetrahedral site; O_h designates the octahedral site. Asterisks designate MAS sidebands.

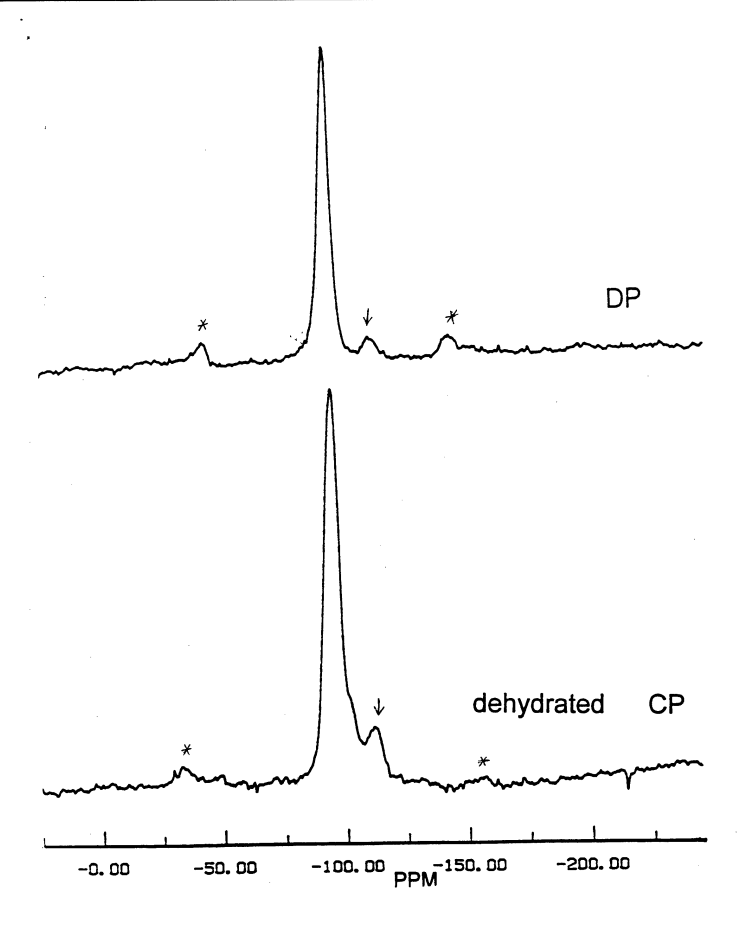


Figure 2. ²⁹Si MAS spectra of Ca-montomorillonite. Upper, DP (direct polarization). Lower, CP (cross polarization). Asterisks denote spinning sidebands. Arrow denotes silica peak.

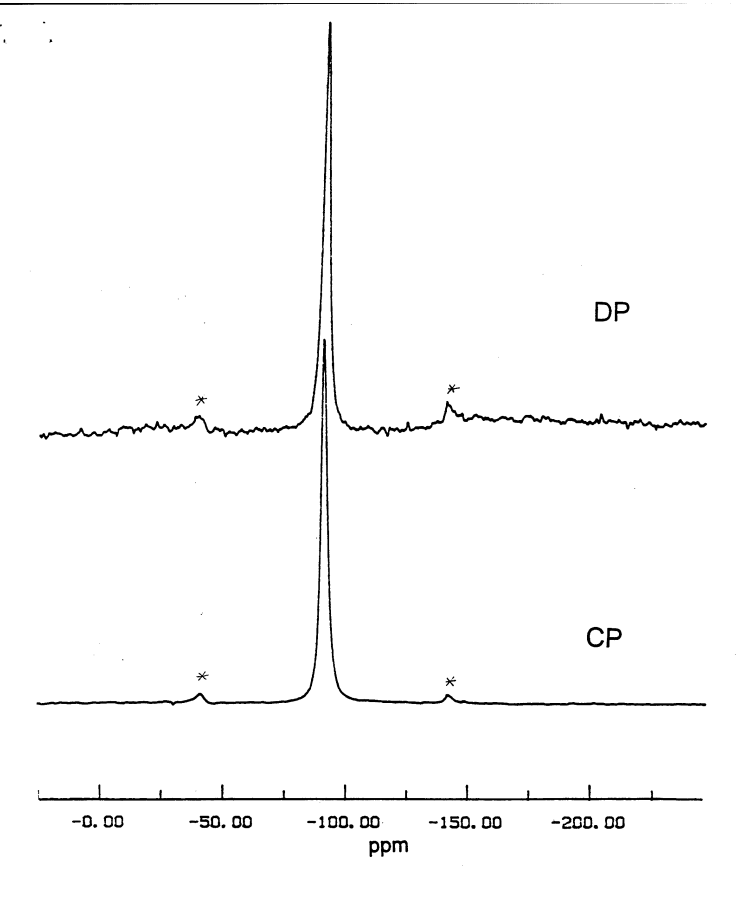


Figure 3. ²⁹Si MAS spectra of hydrated kaolinite. Asterisks denote MAS sidebands.

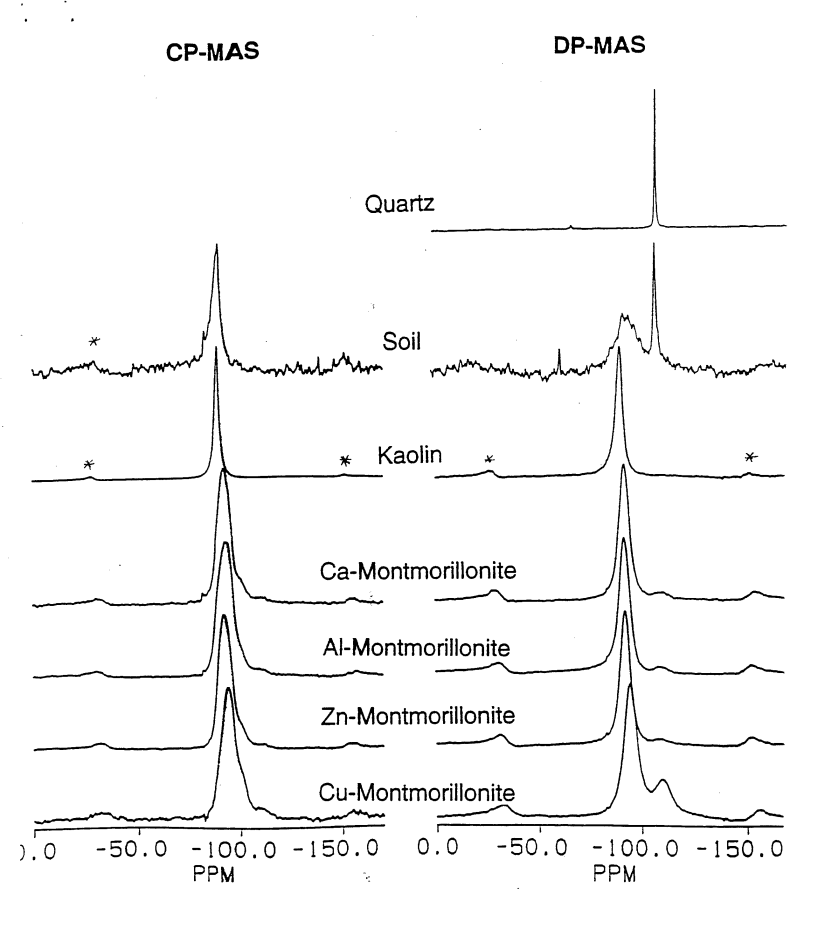


Figure 4. ²⁹Si MAS spectra of clays and soils. Asterisks denote MAS sidebands.

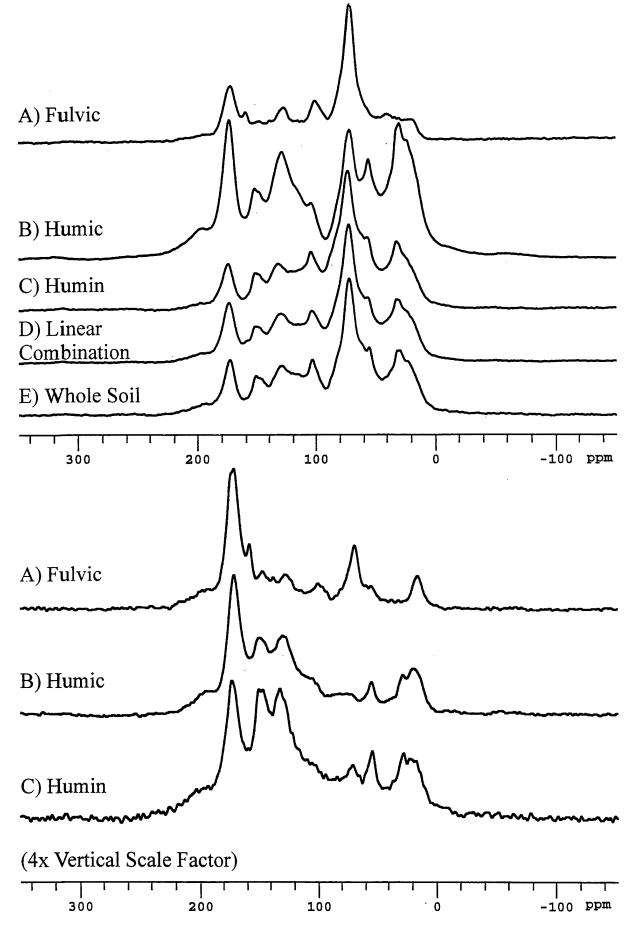


Figure 5. ¹³C CP-MAS spectra of A) fulvic acid, B) humic acid, C) humin from the Uncompandere National Forest and E) whole soil. D) Computergenerated linear combination spectrum: (0.090)(1A) + (0.15)(1B) + (0.76)(1C).

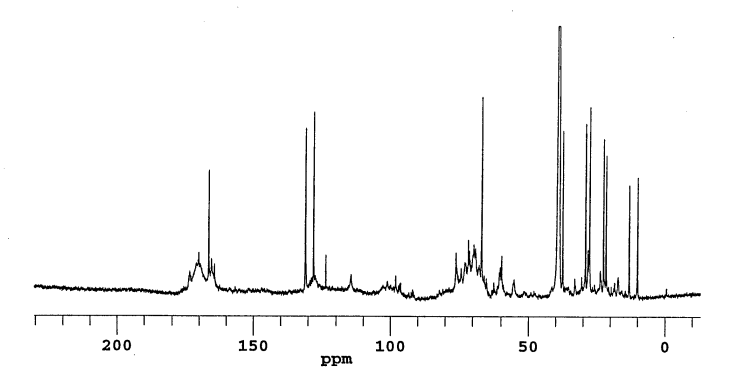


Figure 6. Liquid-solution ^{13}C NMR spectra of the Uncompanger fulvic acid (in $(D_3C)_2SO$).

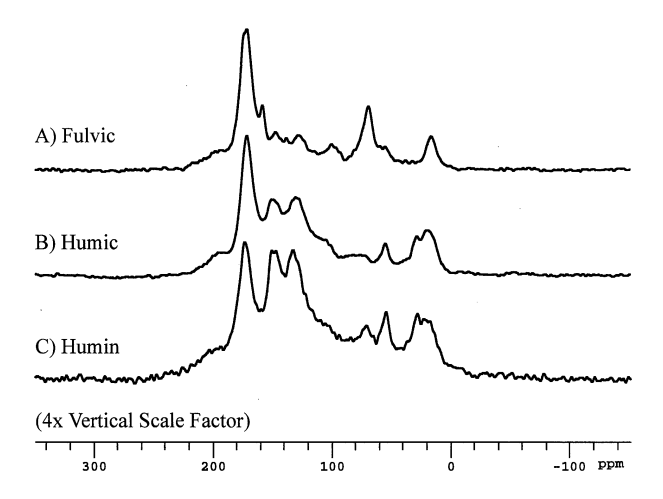


Figure 7. Dipolar-dephasing ¹³C CP-MAS spectra of Uncompanger fulvic acid (A), humic acid (B) and humin (C). 75 µs dipolar-dephasing period.

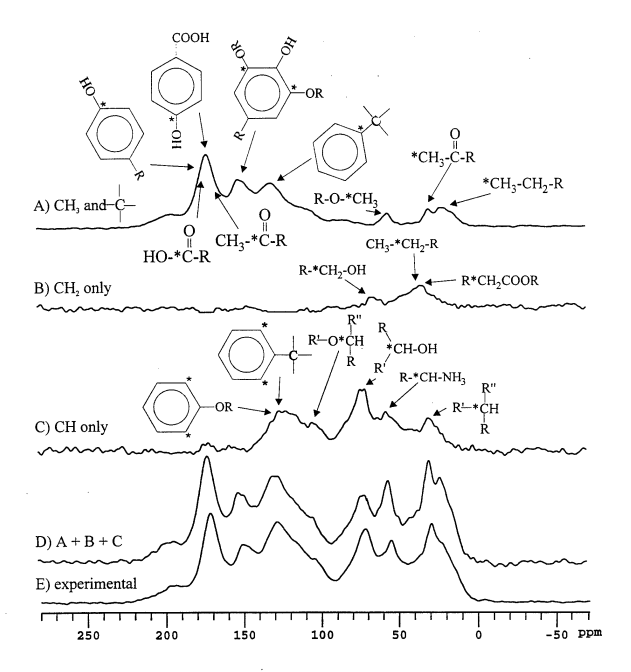
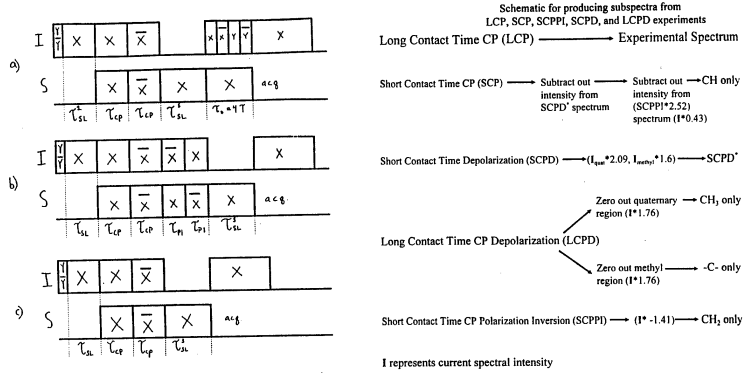


Figure 8. ¹³C CP-MAS spectral editing of the Uncompangre humic acid A) - C), individual component spectra derived according to the published method; D) Composite spectrum obtained from the individual components; E) Experimental CP-MAS spectrum (Fig. 1B).



- a) Standard CP-MAS experiment combined with a depolarization time τ_D . This is the pulse program which produces both SCPD and LCPD spectra (short and long contact time experiments with dephasing).
- b) Standard CP-MAS experiment combined with a polarization inversion period τ_{PI}, which forms the SCPPI spectra.
- c) Standard CP-MAS experiment which is used for both LCP and SCP spectra (long and short contact times).

X. Wu, S. Burns, and K. Zilm, J. Magn. Reson. A 111, 29 (1994)

Figure 9. General procedure used for obtaining CP-based spectral editing of ¹³C NMR spectra of the type shown in Figure 8.

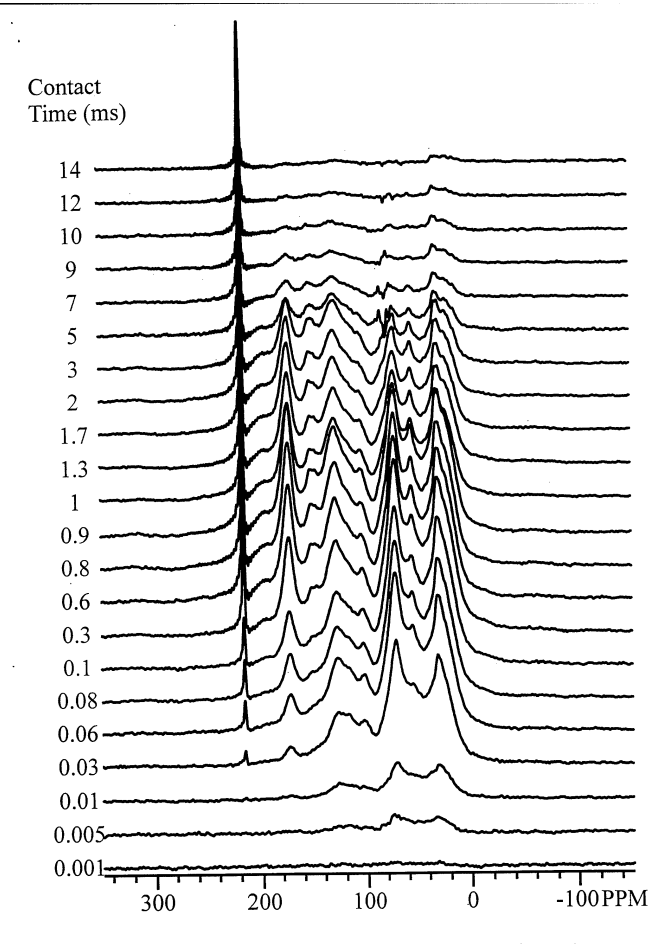


Figure 10. ¹³C spectra of a variable contact-time CP-MAS experiment on Uncompangre humic acid.

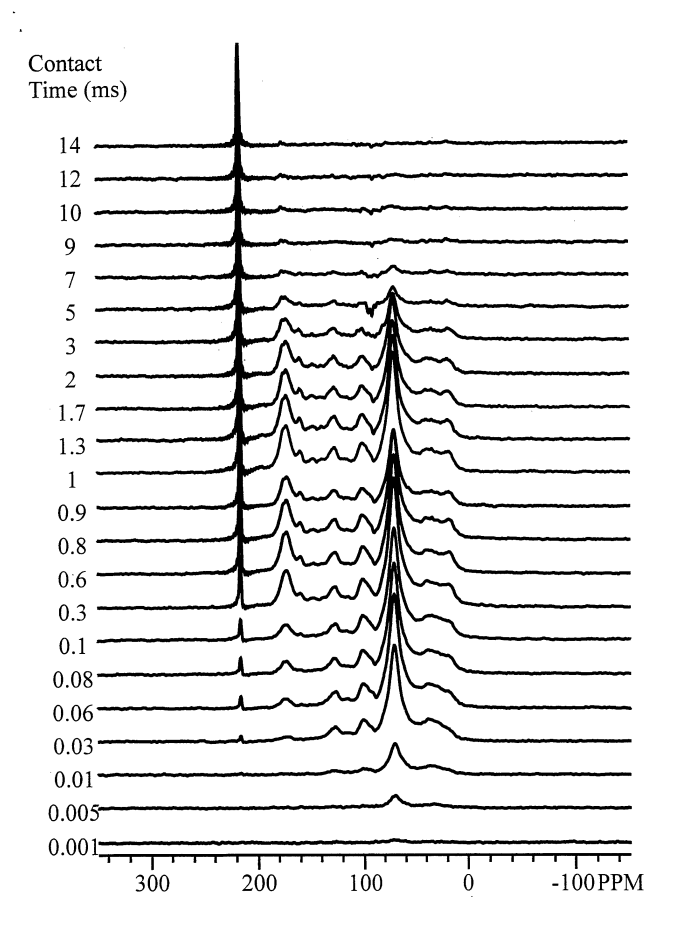


Figure 11. ¹³C spectra of a variable contact-time CP-MAS experiment on Uncompangre fulvic acid.

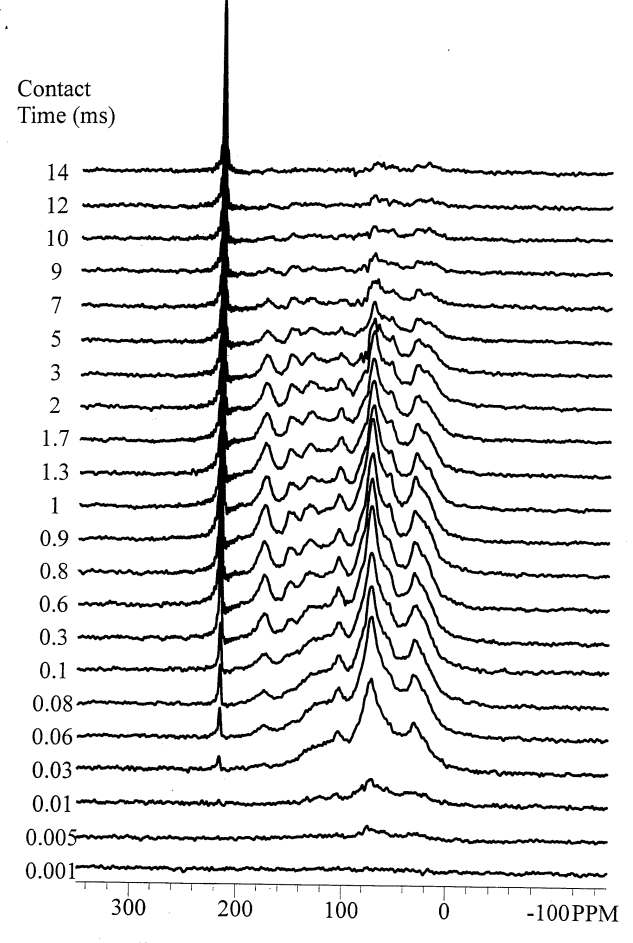


Figure 12. ¹³C spectra of a variable contact-time CP-MAS experiment on Uncompangre humin.

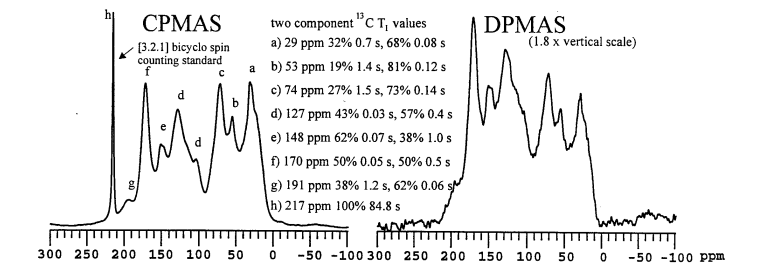
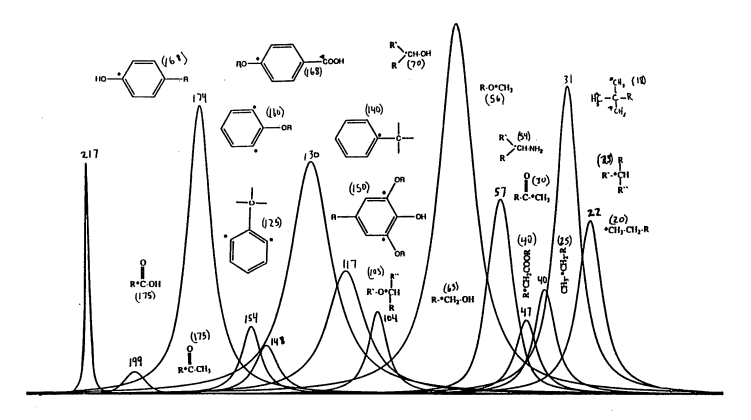


Figure 13. Typical high S/N ¹³C CP-MAS (A) and DP-MAS (B) spectra of Uncompange humic acid, used for ¹³C spin counting. Intensity reference compound in the sample of A.



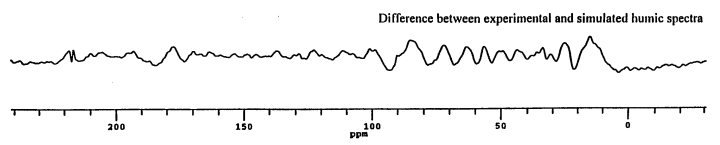


Figure 14. Spectral deconvolution of the ¹³C CP-MAS spectrum of Uncompangre humic acid.

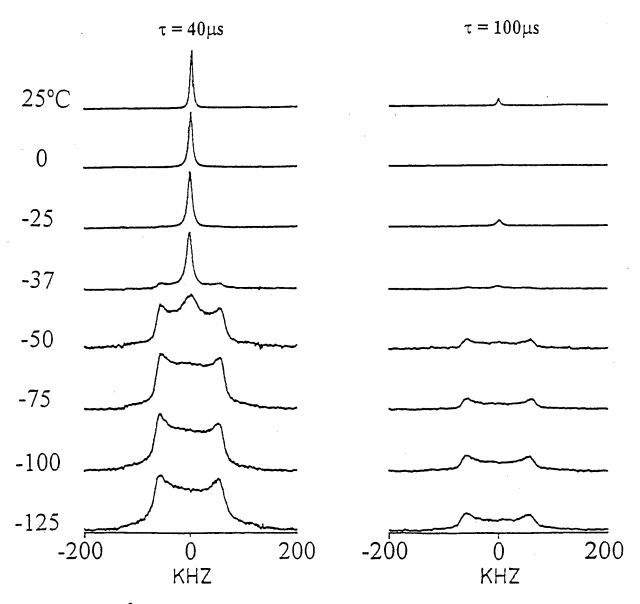


Figure 15. 2 H NMR spectra of HOCD $_2$ CD $_2$ OH adsorbed on Camontmorillonite as a function of temperature and quadrupole echo period (τ), as indicated.

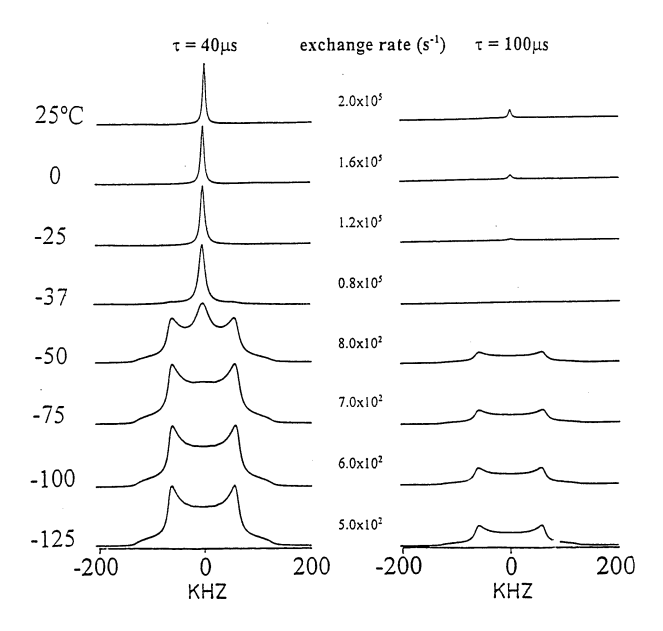


Figure 16. Computer simulations of the ²H NMR spectra shown in Fig. 15.

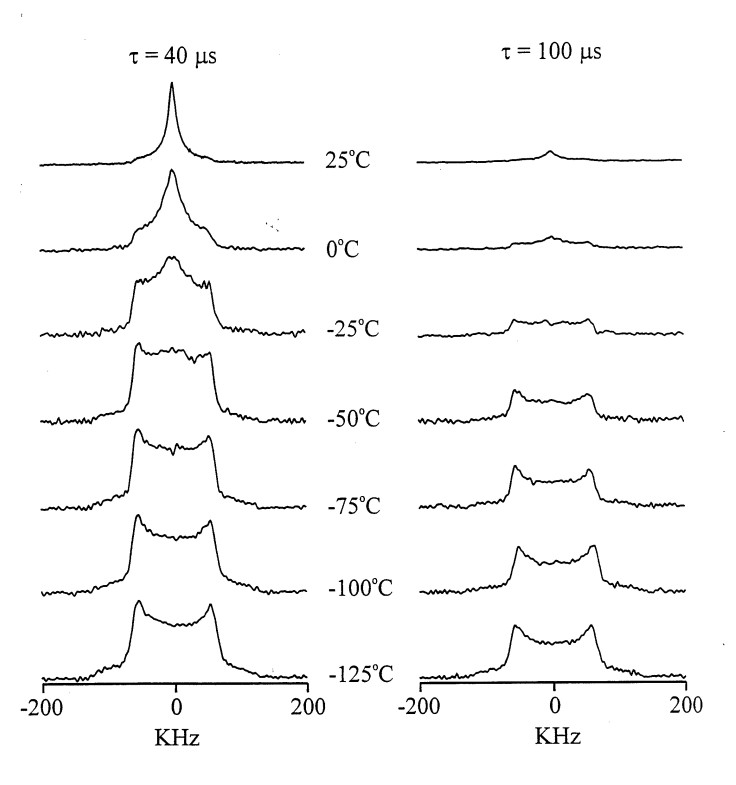


Figure 17. ²H NMR spectra of a sample of HOCD₂CD₂OH adsorbed on dry Uncompange humic acid as a function of temperature and quadrupole echo delay period (τ), as indicated.

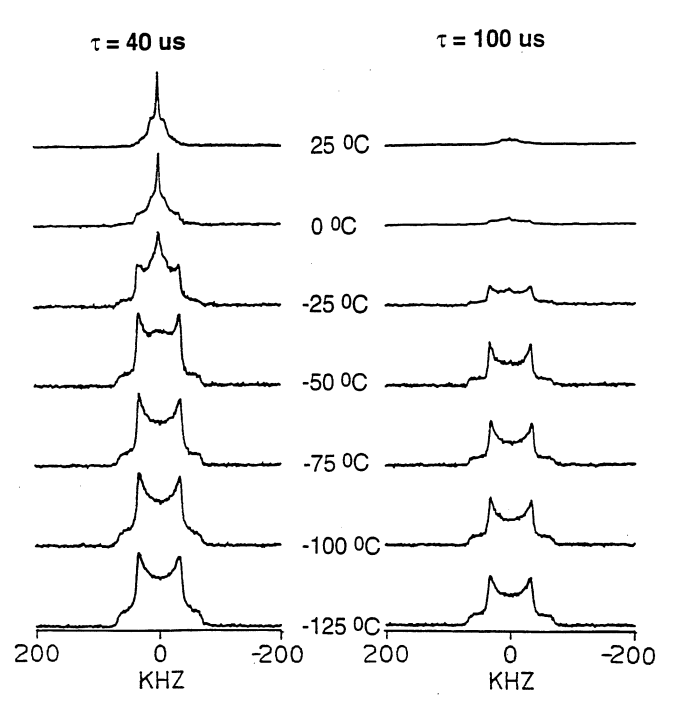


Figure 18. 2 H NMR spectra of a sample of C_6D_6 adsorbed on Camontmorillonite (loading level: 1.2 % wt), as a function of temperature and quadrupole echo delay period (τ), as indicated.

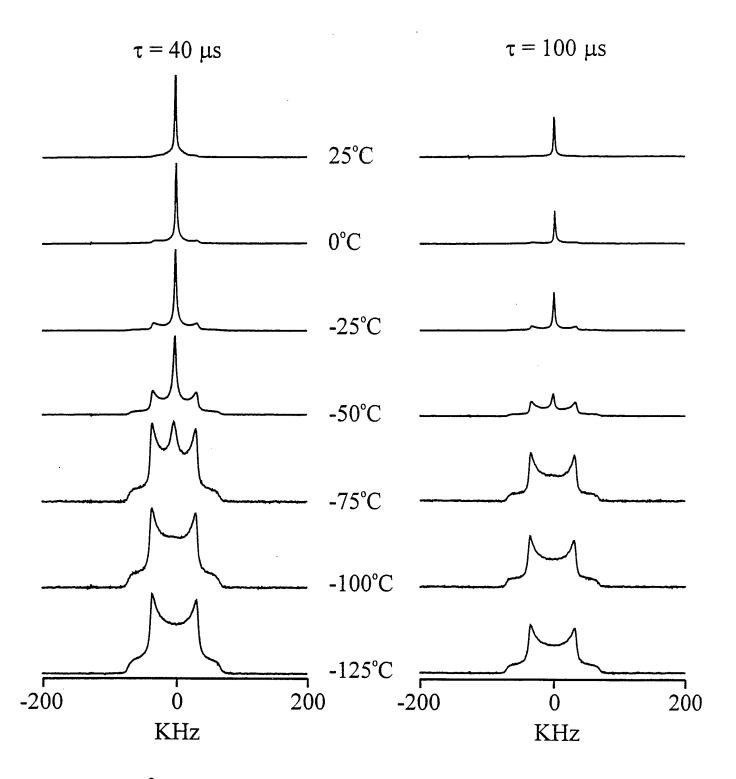


Figure 19. 2 H NMR spectra of a sample of C_6D_6 adsorbed on Camontmorillonite (loading level: 3.9 % wt), as a function of temperature and quadrupole echo delay period (τ) , as indicated.

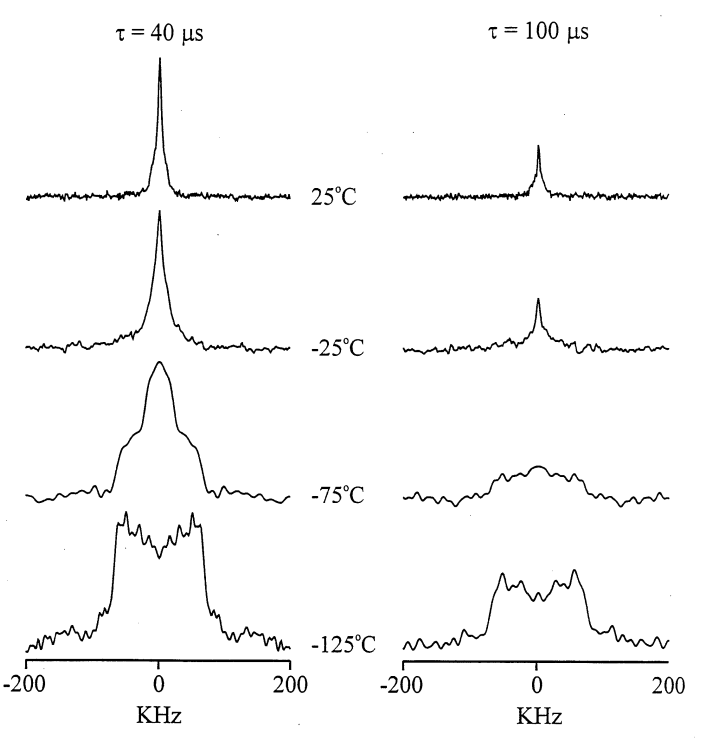


Figure 20. 2 H NMR spectra of a sample of $Cl_2C=CDCl$ adsorbed on Camontmorillonite, as a function of temperature and quadrupole echo delay period (τ), as indicated.

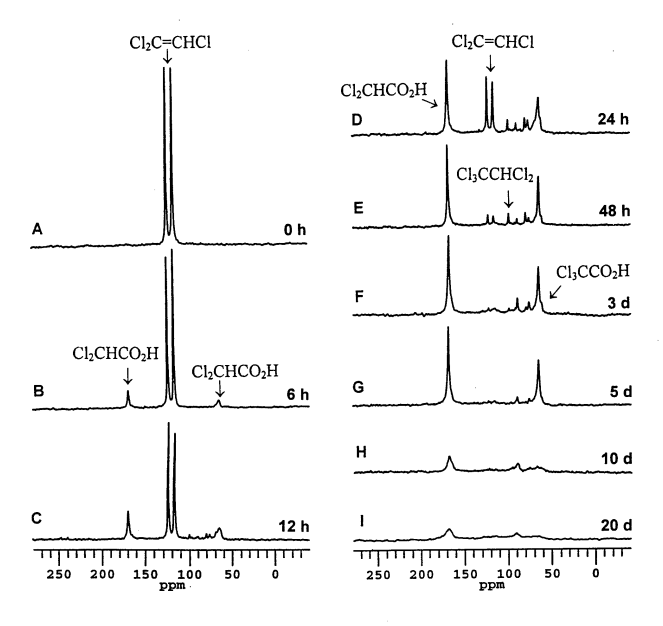


Figure 21. 13 C DP-MAS spectra of Cl_2C =CHCl adsorbed on Camontmorillonite, subjected to near-UV radiation for various periods of time, as indicated (h = hours, d = days).

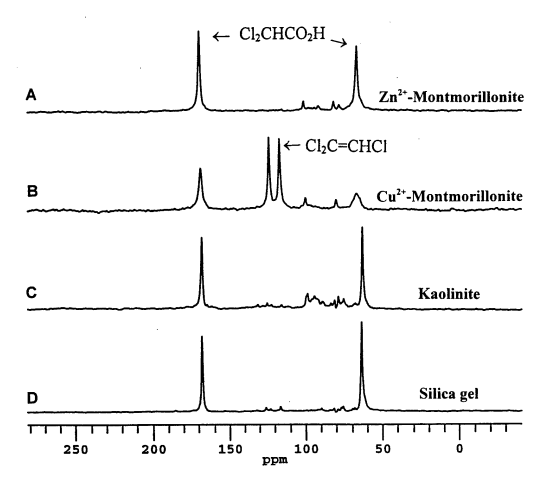


Figure 22. ¹³C DP-MAS spectra of Cl₂C=CHCl adsorbed on the indicated substrates, subjected to near-UV radiation for 48 hours.

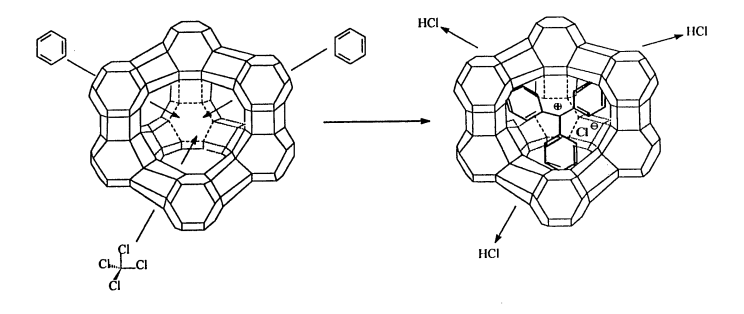


Figure 23. Cartoon representation of the "ship-in-a-bottle" synthesis of $(C_6H_5)_3C^+$ ions inside the cavity of HY zeolite via Friedel-Crafts processes.

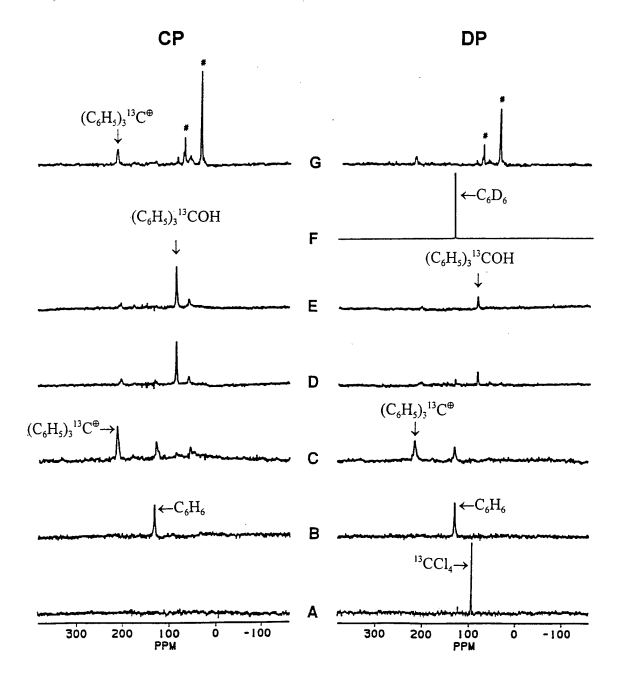


Figure 24. ¹³C DP-MAS and CP-MAS NMR spectra of the reaction product of ¹³CCl₄ with benzene in HY, and related samples. (A) ¹³CCl₄ mixed with preheated HY; (B) benzene mixed with preheated HY; (C) ¹³CCl₄ and benzene mixed with preheated HY; (D) sample 24C exposed to ambient moisture; (E) solid residue of sample 24D washed with C₆D₆; (F) liquid extract from C₆D₆ extraction of sample 24D; (G) product of reaction of sample 24D with LiAl[OC(CH₃)₃]₃H (peaks marked with # at top correspond to the signals from the reducing agent).

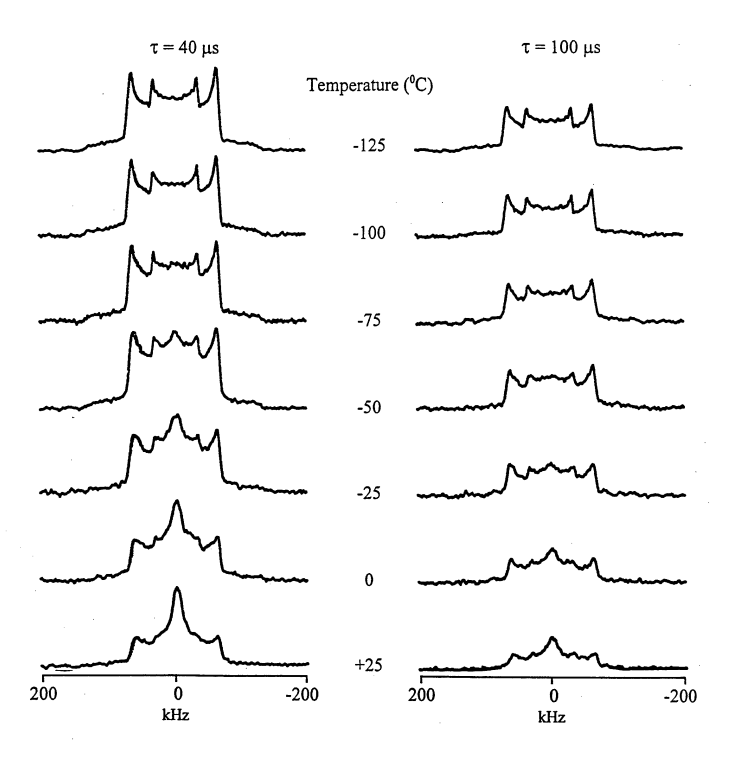


Figure 25. 2 H NMR spectra of $(C_6D_5)_3C^+$ adsorbed primarily in the HY zeolite cavities, as a function of temperature and quadrupole echo delay period (τ) , as indicated.

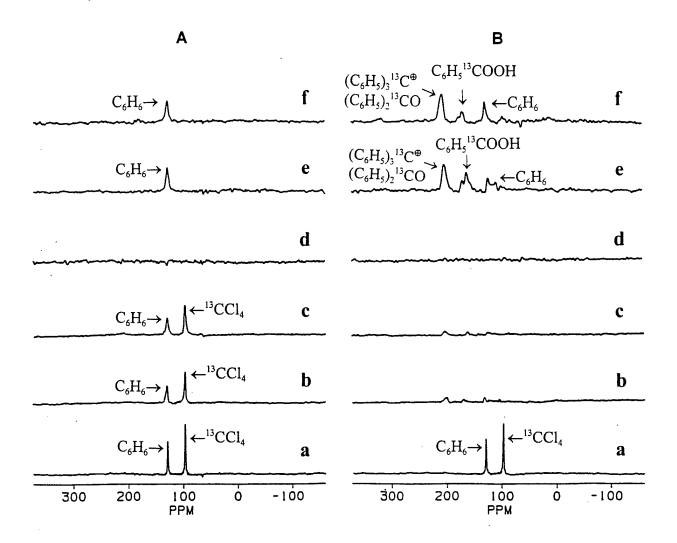


Figure 26. ¹³C DP-MAS (a,b,c) and CP-MAS (d,e,f) NMR spectra of samples of benzene and ¹³C-labeled carbon tetrachloride on (A-a, A-d) kaolinite, (A-b, A-e) Ca-montmorillonite, (A-c, A-f) K-10 montmorillonite, (B-a, B-d) Zn²⁺-exchanged kaolinite, (B-b, B-e) Zn²⁺-exchanged Camontmorillonite and (B-c, B-f) Zn²⁺-exchanged K-10 montmorillonite.

Final Invention Report

AFOSR Grant F49620-95-1-0192 Organic

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P.I. Gary E. Maciel

Co-P.I. Willard L. Lindsay

There were no inventions.

Gary E. Maciel

4/20/98

Date